

Flex Databases Platform

Unified eClinical platform for CRO, biotech, and pharma

GENERAL
CTMS
PM & BUJGETING
COMPLIANCE & SAFETY

Image: Project Management & Budgeting
Image: Project Management & Budgeting
Image: Project Management & Budgeting
Learning Management & System

Image: Project Management & Budgeting
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About Flex Databases

We develop a comprehensive clinical trial management solution since 2011.

What sets us apart

Easy implementation
Full compliance
Complete data safety
Flexible solution

- 10+ years on the market
- 4 offices in the US, Europe, and Asia
- All-in-one platform from study planning to safety database
- System implementation in 3 to 10 weeks
- A robust backup & disaster recovery and data protection strategy, including distributed data storages around the world
- All major international and local regulations covered, including FDA 21 CFR Part 11, GDPR, etc.
- Reports, workflows, trackers are under user configuration. Web-based solution, no installation required

Some of our clients

Leading CROs, pharma, and biotech











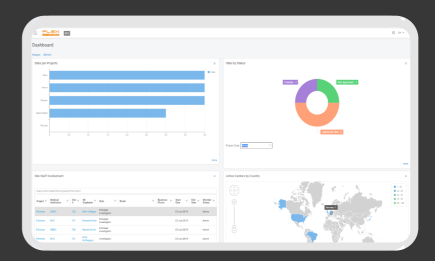






Investigators & Sites Management

Module Overview



Organize all the information on investigators, medical institutions, sites, regulatory authorities, and vendors

Sites statuses, addresses, documents, contracts, qualification, vendors, submissions, site staff, timelines and so much more at hand.

Find all the related information in few clicks

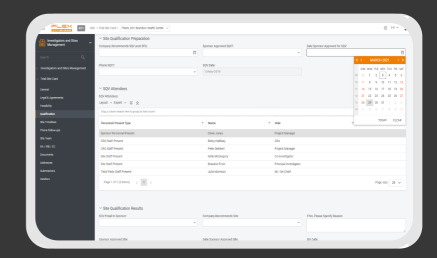
Keep all your Medical Institutions, Sites, and Sites Staff information in a clear organized manner.

Manage submissions packages across countries, studies and sites in one interface.



Investigators & Sites Management

List of Features



Regulators and investigators

- RA, IRB, and EC information: address, contact person, periodicity of session, terms of submission, and other key information
- Investigators information: contacts,
 CVs, certificates, and licenses,
 experience, studies, therapeutic areas,
 etc.

Site management

- Sites information: address, accreditation certificates, study team, and studies
- Site team assignment
- Site performance
- Capturing and tracking of site communication

Hospitals, clinics, and vendors

- Hospitals and out-patient clinics (addresses, description, equipment, capacity)
- Vendor information (address, pricing, services)

Tracking and reporting

- Documents tracking: contracts, site regulatory documents, licenses, and certificates
- Centralized IRB/LEC submissions and approval tracking
- Ad-hoc reporting tool for crossproject and cross-module reporting (graphs, widgets)